

11-24-00

01

11/22/00  
 Jc962 U.S. PTO

# UTILITY PATENT APPLICATION TRANSMITTAL

(Only for new nonprovisional applications under 37 CFR 1.53(b))

Attorney Docket No. 09792909-4673

First Named Inventor or Application Identifier

Tsuyonobu Hatazawa

Express Mail Label No: EL654782576US

Jc916 U.S. PTO  
 09/18/00  
 11/22/00

ADDRESS TO: Assistant Commissioner for Patents  
 Box Patent Application  
 Washington, DC 20231

## APPLICATION ELEMENTS

See MPEP chapter 600 concerning utility patent application contents.

1. ☒ Specification Total Pages 37
2. ☒ Drawing(s) (35USC 113) Total Pages 6
3. ☒ Declaration and Power of Attorney Total Pages 5
  - a. ☒ Unexecuted(original or copy)
  - b. ☐ Copy from prior application (37CFR 1.63(d))  
 (for continuation/divisional with Box 14 completed)  
 [Note Box 4 Below]  
 i. ☐ **DELETION OF INVENTOR(S)**  
 Signed statement attached deleting Inventor(s) named in the prior application, see 37 CFR 1.63(d)(2) and 1.33(b).
4. ☐ Incorporation By Reference (usable if Box 3b is checked)  
 The entire disclosure of the prior application, from which  
 a copy of the oath or declaration is supplied under Box 3b, is considered as being part of the disclosure of the accompanying application and is hereby incorporated by reference therein.

## ACCOMPANYING APPLICATION PARTS

5. ☐ Assignment Papers (cover sheet & documentation)
6. ☒ Letter under 37 CFR 1.41(c).
7. ☐ English Translation Document (if applicable)
8. ☐ Information Disclosure Statement (IDS)/PTO-1449 ☐ Copies of IDS Citations
9. ☒ Preliminary Amendment
10. ☒ Return Receipt Postcard (MPEP 503)  
 (Should be specifically itemized)
11. ☐ Small Entity ☐ Statement filed in prior Application, Status still proper and desired
12. ☒ Certified copy of Japanese priority application No. P11-341373 filed November 30, 1999

14. If a CONTINUING APPLICATION, check appropriate box and supply the requisite information:

☐ Continuation ☐ Divisional ☐ Continuation-in-part (CIP) ☐ of prior application No:

## CLAIMS AS FILED

	(1) FOR	(2) NUMBER FILED	(3) NUMBER EXTRA	(4) RATE	(5) BASIC FEE \$710.00
TOTAL CLAIMS	20	11		18.00	8.00
INDEPENDENT CLAIMS	03	1		80.00	0.00
		ANY MULTIPLE DEPENDENT CLAIMS? <input type="checkbox"/> YES <input checked="" type="checkbox"/> NO			0.00
		TOTAL FEE			\$710.00

☒ The Commissioner is hereby authorized to charge any additional fees which may be required in connection with this application, or credit any overpayment to ACCOUNT NO. 19-3140. A duplicate copy of this sheet is enclosed.

☒ A check in the amount of \$710.00 to cover the filing fee is enclosed.

15. CORRESPONDENCE ADDRESS: David R. Metzger  
 SONNENSCHN NATH & ROSENTHAL  
 P.O. Box #061080  
 Wacker Drive Station, Sears Tower  
 Chicago, IL 60606-1080  
 Customer ID #26263  
 Telephone (312) 876-8000 - Fax (312) 876-7934

SIGNATURE

*David R. Metzger*  
 David R. Metzger (Reg. No. 32,919)

Date: November 22, 2000

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

**PRELIMINARY AMENDMENT ACCOMPANYING APPLICATION**

APPLICANT: Tsuyonobu Hatazawa, et al. ATTY. DOCKET NO. 09792909-4673

SERIAL NO.

DATE FILED:

INVENTION: "NONAQUEOUS ELECTROLYTE BATTERY"

Assistant Commissioner of Patents  
Washington, D.C. 20231

S I R:

Between the title and the heading "Background of the Invention" on page 1, insert the following:

--RELATED APPLICATION DATA

The present application claims priority to Japanese Application No. P11-341373 filed November 30, 1999, which application is incorporated herein by reference to the extent permitted by law.

Respectfully submitted,



(Reg. No. 32,919)

David R. Metzger  
SONNENSCHN NATH & ROSENTHAL  
P.O. Box #061080  
Wacker Drive Station  
Sears Tower  
Chicago, IL 60606-1080  
Customer #26263

Attorneys for Applicant(s)

## NONAQUEOUS ELECTROLYTE BATTERY

### BACKGROUND OF THE INVENTION

The present invention relates to a nonaqueous electrolyte battery in which a battery element is contained in an outer covering member composed of a laminated film.

In recent years, along with strong demands toward reductions in size and weight in the field of portable electronic equipment such as a camera integrated VTR, a cellular phone, and a portable computer, batteries used as portable power sources of the portable electronic equipment have been also required to be reduced in size and weight. To meet such a requirement, studies have been extensively made to develop thin type or foldable type batteries, particularly, secondary batteries, more particularly, nonaqueous electrolyte batteries (so-called lithium batteries).

As an electrolyte used for the above-described foldable battery, a solid electrolyte has been actively made. In particular, a gel electrolyte composed of a solid electrolyte containing a plasticizer and a high polymer solid electrolyte including a lithium salt dissolved in a high polymer have become a focus of

attention.

On the other hand, to attain the reductions in size and weight of batteries, studies have been also made to develop batteries of a type including a battery element enclosed in a plastic film or a so-called laminated film obtained by laminating a metal film to a plastic film.

The so-called laminated film, however, has a problem. For a conventional battery using a metal container, the metal container having a high rigidity is little deformed even if a pressure in the battery is raised. On the contrary, for a battery using the laminated film, if the battery is erroneously charged or discharged due to a failure of a control circuit or the like or is left under an abnormal high-temperature environment, then the laminated film having a low rigidity is possibly swelled by an increase in pressure in the battery due to gas occurred by the abnormal reaction of constituent materials of the battery.

#### SUMMARY OF THE INVENTION

An object of the present invention is to provide a nonaqueous electrolyte battery capable of preventing the swelling of an outer covering member composed of a laminated film, even if gas occurs in a battery element

of the battery.

To achieve the above object, according to the present invention, there is provided a nonaqueous electrolyte battery including a battery element contained in an outer covering member composed of a laminated film and sealed therein by heat seal, and a gas absorbable material interposed between the outermost layer of said outer covering member and said battery element.

With this configuration, since the gas absorbable material is interposed between the outermost layer of the outer covering member and the battery element, even if gas occurs in the battery element when the battery undergoes an abnormal environmental change, for example, an abnormal temperature rise, the gas is absorbed in the gas absorbable material, with a result that the outer covering member is prevented from being swelled by the gas.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is an exploded perspective view of a nonaqueous electrolyte battery according to the present invention, showing a state before a battery element is assembled in an outer covering;

Fig. 2 is a schematic perspective view of the

nonaqueous electrolyte battery shown in Fig. 1, showing a state after the battery element is assembled in the outer covering;

Fig. 3 is a schematic perspective view showing a state in which the entire surface of the battery element is covered with a gas absorbable member;

Fig. 4 is front, top, and side views showing volume loss portions in the state shown in Fig. 3 in which the entire surface of the battery element is covered with a gas absorbable member;

Fig. 5 is a schematic perspective view showing a state in which gas absorbable members previously formed by molding are attached to the battery element;

Fig. 6 is a schematic perspective view showing a state in which a gas absorbable member is formed in a space between the outer covering member and the battery element by filling the space with a resin mixture containing a gas absorbable material;

Fig. 7 is a sectional view showing a laminated film having a gas absorbable layer;

Fig. 8 is a sectional view showing another laminated film having a gas absorbable layer;

Fig. 9 is a schematic perspective view showing dimensions of a winding type battery element produced in

Examples 1, 2 and 3; and

Fig. 10 is a schematic perspective view showing dimensions of a flat type battery element produced in Examples 4, 5 and 6.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

Hereinafter, an embodiment of a nonaqueous electrolyte battery of the present invention will be described with reference to the drawings.

A nonaqueous electrolyte battery of the present invention is represented by a solid electrolyte battery or a gel electrolyte battery having a configuration shown in Figs. 1 and 2. Referring to these figures, a battery element 1 including a solid electrolyte or a gel electrolyte interposed between a positive active material and a negative active material is contained in an outer covering member 2, typically, a laminated film and is sealed in the outer covering member 2 with its periphery bonded thereto by heat seal.

The battery element 1 has a negative electrode terminal lead 3 electrically connected to a negative electrode of the battery element 1, and a positive electrode terminal lead 4 electrically connected to a positive electrode of the battery element 1. The negative

and positive electrode terminal leads 3 and 4 are led out of the outer covering member 2.

In the nonaqueous electrolyte battery according to this embodiment, gas absorbable members 5 and 6 are inserted between the outermost layer of the laminated film as the outer covering member 2 and at least one or more planes of the battery element 1. In the example shown in Figs. 1 and 2, the battery element 1 is of a winding type, and the above-described at least one or more planes of the battery element 1 are specified by two winding end surfaces of the battery element 1. With the insertion of these gas absorbable members 5 and 6, if gas occurs in the battery element 1 due to an abnormal environmental change such as an abnormal temperature rise or overcharging, the gas is absorbed in the gas absorbable members 5 and 6, with a result that the battery is prevented from being deformed by swelling of the outer covering member 2. Each of the gas absorbable members 5 and 6 is made from a gas absorbable material and a resin material.

The gas absorbable material is represented by a porous metal compound or porous carbon material. Examples of the porous metal compounds may include zeolite, alumina, molecular sieve, titania, silica gel, and



zirconia, and examples of the porous carbon materials may include activated carbon, and carbon molecular sieve.

Other than the above-described porous metal compound or porous carbon material, a metal compound selected from magnesium sulfate, calcium chloride, and phosphorus pentaoxide, or a metal selected from Ni, Pt, Pd, Ca, Sr, Ba, Ti, Zr, Hf, V, Nb, Ta, Cr, Mo, W, and Fe may be used as the gas absorbable material.

The above-described gas absorbable materials may be used singly or in combination.

Since the components and amount of gas occurring in the battery element 1 are dependent on the kinds and amounts of an electrolyte salt and an electrolyte solvent used for a gel electrolyte layer, the kind and amount of a gas absorbable material to be inserted are largely dependent on a configuration of the electrolyte and the reaction amount thereof.

The selection of the kind and amount of a gas absorbable material depending on the kind of an electrolyte and the reaction amount thereof was examined by way of the following experiment.

A sample battery of a 500 mh grade, using  $\text{LiCoO}_2$  as a positive active material and graphite as a negative active material, and using 1.5 g of an electrolyte

03754400

solution, was produced. In this experiment, the electrolyte solution was prepared by dissolving  $\text{LiPF}_6$  as an electrolyte salt in a propylene carbonate as an electrolyte solvent exhibiting a high reactivity with graphite at a concentration of 1.0 mol/kg. After the sample battery overcharged at 4.35 V was left at  $100^\circ\text{C}$  for 100 hr, gas occurred in the sample battery was analyzed using a GC-MS (Gas Chromatography-Mass Spectrometer). As a result, it was found that the reacted propylene carbonate was about 10% of the total amount inserted and that the gas mainly contained carbon dioxide, propane, propylene, and evaporated unreacted propylene carbonate.

After the sample battery was cooled to ordinary temperature, the gas remaining in the sample battery was also analyzed. As a result, it was found that carbon dioxide, propane, and propylene produced by reaction of slightly less than 10% of 1.5 g of the propylene carbonate electrolyte solution was present as the gas components. In this experiment, it was proved that the above gas components produced by the reaction (or decomposition) of about 10% of the electrolyte solution are nearly perfectly absorbable in 0.07 g of 5A carbon molecular sieve having an absorption surface area of about  $3000 \text{ m}^2/\text{g}$ . This means that, in the case of using

0.7 g of 5A carbon molecular sieve as the gas absorbable material, even if the propylene carbonate electrolyte solution is nearly 100% decomposed, gas components produced by such decomposition can be absorbed in the gas absorbable material.

In the above experiment, the sample battery is produced by the combination of the materials having high reactivity with each other; however, in actual, a battery is produced by combination of materials having low reactivity with each other. Accordingly, for an actual battery, the inserted amount of a gas absorbable material may become smaller than the above-described experimentally determined amount of 5A carbon molecular sieve used as the gas absorbable material.

The optimum kind and amount of the gas absorbable material, which are largely dependent on the combination of constituent materials of a battery, should be selected for each battery from the practical viewpoint.

The resin material used for each of the gas absorbable members 5 and 6 may be at least one kind or more selected from polyethylene, polypropylene, polyethylene terephthalate, polyimide, polyamideimide, polyamide, polyphenylene sulfide, tetrafluoroethylene-perfluoroalkylvinylether copolymer, polyvinylidene

fluoride, polytetrafluoroethylene, ethylene fluoride-polypropylene copolymer, ethylene-tetrafluoroethylene copolymer, polyvinyl fluoride, epoxy resin, acrylic resin, and methacrylic resin.

The resin material used for each of the gas absorbable members 5 and 6 may be also at least one kind or more cross-linked resins. Examples of monomers for the cross-linked resins may include 2-hydroxyethyl (meth)acrylate, 2-hydroxypropyl (meth)acrylate, glycerol (meth)acrylate, 2-acrylamide-2-methylpropanesulfonic acid, sodium sulfonate-ethyl(meth)acrylate, (meth)acrylamide, N,N-dimethylaminoethyl(meth)acrylate, N,N-diethylaminoethyl(meth)acrylate, N-isopropylacrylamide, N,N-dimethylaminopropylacrylamide, (meth)acrylic acid, sodium (meth)acrylate, 2-(meth)acryloyloxyethylsuccinic acid, 2-(meth)acryloyloxyethyl phthalate, 2-(meth)acryloyloxyethyl hexahydrophthalate,  $\omega$ -carboxypolycaprolactonemonoacrylate, EO modified phosphoric (meth)acrylate, polyethyleneglycol (meth)acrylate, acryloylmorpholine, p-styrenesulfonic acid, vinylsulfonic acid, allylsulfonic acid, (meth)acrylic ethylsulfonic acid, (poly)ethyleneglycol di(meth)acrylate (for example, produced by Shin-Nakamura Chemical Co., Ltd. under the trade names of NK Ester A-200, A-400, A-600, A-1000, 4G,

9G, 14G, and 23G; and produced by Kyoeisha Chemical Co., Ltd. under the trade names of Light Ester 4EG, 9EG, 14EG, 9EG-A, and 14EG-A), (poly)ethyleneglycol mono(meth)acrylate (for example, produced by Shin-Nakamura Chemical Co., Ltd. under the trade names of NK Ester AMP-10G, AMP-20G, AMP-60G, and AMP-90G; and produced by Kyoeisha Chemical Co., Ltd. under the trade names of Light Ester MC, 130MA, and 041MA), and ethoxyeated polypropyleneglycol dimethacrylate (produced by Shin-Nakamura Chemical Co., Ltd. under the trade name of NK Ester 1206PE). These materials may be hardened by irradiation of heat, ultraviolet rays, visual light, electron rays, or radiation thereto.

In the case of adopting the hardening process by irradiation of ultraviolet rays, a material activated by ultraviolet rays may be used as an initiator. Examples of such initiators may include a sulfide such as sodium methyldithiocarbamate sulfide, tetramethylthiuram monosulfide, diphenyl monosulfide, dibenzothiazoyl monosulfide, or disulfide; thioxanthone, and a derivative thereof such as ethylthioxanthone, 2-chlorothioxanthone, diethylthioxanthone, or diisopropylthioxanthone; a diazo compound such as hydrazone, azoisobutyronitrile, or benzenediazonium; an aromatic carbonyl compound such as

benzoin, benzoinmethylether, benzoinethylether,  
benzoinisopropylether, benzophenone,  
dimethylaminobenzophenone, Michler's ketone,  
benzilantraquinone, t-butylantraquinone, 2-  
methylantraquinone, 2-ethylantraquinone, 2-  
aminoantraquinone, 2-chloroantraquinone, benzildimethyl  
ketal, or methylphenyl glyoxylate; an acetophenone  
derivative such as 1-hydroxycyclohexyl phenyl ketone, 2-  
hydroxy-2-methyl-1-phenylpropane-1-on, 4-(2-  
hydroxyethoxy)phenyl(2-hydroxy-2-propyl) ketone, 2,2-  
diethoxyacetophenone, or 2,2-dimethoxyacetophenone; a  
dialkylamino benzoate such as 4-dimethylaminomethyl  
zenboate, 4-dimethylaminoethyl zenboate, 4-  
dimethylaminobutyl zenboate, or 4-dimethylaminoisopropyl  
zenboate; a peroxide such as benzoil peroxide, di-t-butyl  
peroxide, dicumyl peroxide, or cumene hydroperoxide; an  
acridine derivative such as 9-phenylacridine, 9-p-  
methoxyphenylaciridine, 9-acethylaminoacridine, or  
benzacridine; a phenazine derivative such as 9,10-  
dimethylzenzphenazine, 9-methylbenzphenazine, or 10-  
methoxybenzphazine; a quinoxaline derivative such as  
4,4',4''-trimethoxy-2,3-diphenylquinoxaline; 2,4,5-  
triphenylimidazolyl dimer; ketone halide; and an acylated  
phosphide such as acylphophine oxide or acylphosphonate.

In the case of adopting the hardening process by irradiation of visual light, a material activated by visual light may be used as an initiator. Examples of such initiators may include 2-nitrofluorene, 2,4,6-tris(trichloromethyl)-1,3,5-triazine, 3-3'-carbonylbiscoumarin, and thio-Michler's ketone.

The above-described initiators may be used singly or in combination.

The added amount of the initiator may be in a range of 0.01 to 20 parts by weight, preferably, in a range of 0.1 to 5 parts by weight. If the added amount of the initiator is less than 0.01 part by weight, the initiator cannot sufficiently exhibit the effect of promoting the crosslinking reaction, with a result that the crosslinking point is lowered and thereby the produced cross-linked resin tends to become softer. Meanwhile, if the added amount of the initiator is more than 20 parts by weight, the initiator excessively promotes the crosslinking reaction, with a result that the produced cross-linked resin tends to become brittle and be changed in color.

In the case of irradiation of ultraviolet rays as activation rays, an extra-high pressure mercury lamp, a high pressure mercury lamp, a metal halide lamp, xenon

lamp, or a low pressure bactericidal lamp may be used as an irradiation source. The irradiation amount of ultraviolet rays, which is dependent on the kind of a monomer, and the added amount of a photochemical polymerization initiator, may be set in a range of 10 mJ/cm<sup>2</sup> to 10 J/cm<sup>2</sup>. In this case, to prevent adverse effect of oxygen, the irradiation of ultraviolet rays may be performed in an inert gas atmosphere or in a state in which the environmental atmosphere is cutoff by using glass or a transparent film.

Each of the gas absorbable members 5 and 6 is obtained by mixing a powder of the above described gas absorbable material with the above-described resin material and hardening the mixture into a suitable shape, for example, a plate-shape. If a thermoplastic resin is selected as the resin material, the resin mixed with the gas absorbable material may be kneaded and extrusion-molded or injection-molded into a desired shape.

The content of the gas absorbable material may be set in a range of about 0.1 wt% to 95 wt% on the basis of the weight of the resin material. If the content of the gas absorbable material is less than 0.1 wt%, the gas absorption function of the gas absorbable material becomes insufficient, and if the content is more than 95



wt%, it becomes difficult to form the gas absorbable member by molding.

In actual, as described above, since the kind and amount of gas occurring in a battery element are largely dependent on the combination of constituent materials of a battery, the amount of the gas absorbable material may be desirable to be suitably selected for each actual battery.

Assuming that the shortest direction of a battery is defined as the thickness direction of the battery, it is important to make the thickness of the battery as thin as possible while keeping constant the capacity of the battery. However, if the gas absorbable member is inserted in the thickness direction between the battery element 1 and the outer covering member, the thickness of the battery is increased. For a battery having a thickness of 3 mm, if the thickness of the battery is increased by 100  $\mu$ m, the volume energy efficiency of the battery is degraded by 3%, and for a battery having a thickness of 0.5 mm, if the thickness of the battery is increased by 100  $\mu$ m, the volume energy efficiency of the battery is degraded by 20%. It should be noted that the relationship between an increase in thickness of the battery and a loss of volume energy efficiency thereof is

actually dependent on the shape of the battery. Now, assuming a battery to have a shape similar to that of a business card and to have a thickness of 3 mm or more, by inserting the plate-shaped gas absorbable members 5 and 6 between the battery element 1 and the laminated film as the outer covering member 2 as shown in Fig. 1, it is possible to ensure an amount gas absorbable material sufficient for gas absorption while suppressing the volume energy efficiency as much as possible.

In the case of using the plate-shaped gas absorbable member, the thickness thereof may be in a range of 50  $\mu\text{m}$  to 1.0 mm, preferably, in a range of 100  $\mu\text{m}$  to 500  $\mu\text{m}$ . If the thickness is more than 1.0 mm, a loss of volume energy density becomes larger, and if the thickness is less than 50  $\mu\text{m}$ , the formation of the gas absorbable member by molding becomes difficult although the amount of the gas absorbable material becomes sufficient.

In this way, in the nonaqueous electrolyte battery of the present invention, in which the battery element having a positive electrode and a negative electrode is enclosed in the laminated film as the outer covering member, the gas absorbable member may be inserted between a certain portion of the battery element and the

outermost layer of the laminated film.

With respect to the arrangement of the gas absorbable member, if the entire surface of the battery element 1 is covered with the gas absorbable member 8 as shown in Fig. 3, it is possible to easily achieve the object of absorbing gas occurred in the battery element 1; however, in this case, the volume loss portions become larger as shown by hatching in Fig. 4, with a result that the rate of the portions other than the battery element 1 occupied in the battery case having a limited volume becomes higher and thereby the volume energy efficiency of the battery is degraded.

Accordingly, the gas absorbable member containing the gas absorbable material according to the present invention may be arranged at a specific location.

The arrangement of the gas absorbable member will be described in detail with reference to Figs. 5 to 8.

As shown in Fig. 5, the thin plate-shaped gas absorbable members 5 and 6 previously formed by molding are attached to the battery element 1, and the battery element 1 to which the members 5 and 6 are attached is enclosed by the laminated film. In an alternative, the battery element 1 is coated with a hardening resin containing the gas absorbable material, followed by

hardening of the resin mixture, and the battery element 1 covered with the gas absorbable material is enclosed by the laminated film.

In the case of using the laminated film having a battery containing recess at a certain location of the film, there may be adopted an arrangement method of containing the battery element in the recess of the laminated film, filling the space between the wall of the recess and the battery element with a hardening material containing the gas absorbable material, hardening the resin mixture, and enclosing the battery by the laminated film. In this case, as shown in Fig. 6, the gas absorbable member 7 is formed around the battery element 1.

The gas absorbable material may be contained in one film layer, positioned in proximity to the battery, of the film layers constituting the laminated film.

Fig. 7 shows a laminated film 10 having one gas absorbable layer. The laminated film 10 has a resin layer 11 having a thickness of about 10  $\mu\text{m}$  made from nylon or polyethylene terephthalate or the like, an aluminum layer 12 having a thickness of 20  $\mu\text{m}$  to 40  $\mu\text{m}$  made from aluminum, and a gas absorbable layer 13 having a thickness of 30  $\mu\text{m}$  to 50  $\mu\text{m}$  made from a resin,

typically, modified cast polypropylene (CPP) in which the gas absorbable material is diffused. These layers 11, 12, and 13 are stacked in this order from the outside in Fig. 7. Fig. 8 shows a variation of the laminated film 10. In this variation, the gas absorbable layer 13 includes a pair of resin layers 13a made from a resin, typically, modified cast polypropylene and a gas absorbable material 13b held between the resin layers 13a.

In the case of providing the gas absorbable layer as one of the layers constituting the laminated film, the thickness thereof may be in a range from 1  $\mu\text{m}$  to 500  $\mu\text{m}$ , preferably, in a range from 5  $\mu\text{m}$  to 100  $\mu\text{m}$ . If the thickness is more than 500  $\mu\text{m}$ , the loss of volume energy density becomes larger, and if the thickness is less than 1  $\mu\text{m}$ , it is difficult to ensure a sufficient amount of the gas absorbable material.

Next, the configuration of the battery element 1 will be described below.

For a solid electrolyte battery or gel electrolyte battery, a high polymer material is used for a high polymer solid electrolyte. Examples of the high polymer materials may include silicon gel, acrylic gel acrylonitrile gel, polyphosphazene modified polymer, polyethylene oxide, polypropylene oxide, and combined

polymers, cross-linked polymers, and modified polymers thereof; and fluorine based polymers such as polyvinylidene fluoride, a vinylidene fluoride-hexafluoropropylene copolymer, vinylidene fluoride-tetrafluoroethylene copolymer, and vinylidene fluoride-trifluoroethylene copolymer, and mixtures thereof.

A solid electrolyte or gel electrolyte stacked on a positive active material layer or negative active material layer is obtained by impregnating the positive active material layer or negative active material layer with a solution containing a high polymer compound, an electrolyte salt, and a solvent (and further, a plasticizer for the gel electrolyte), and removing the solvent to solidify the high polymer compound and electrolyte salt. Accordingly, the solid electrolyte or gel electrolyte stacked on the positive active material layer or negative active material layer is partially impregnated in the positive active material layer or negative active material layer. In the case of using a cross-linking high polymer compound, after the removal of the solvent, the high polymer compound is cross-linked by light or heat.

The gel electrolyte is composed of a plasticizer containing a lithium salt, and a matrix high polymer in

an amount of 2 wt% to 30 wt%. In this case, an ester, an ether, and a carbonate may be used singly or in combination as one component of the plasticizer.

In preparation of the gel electrolyte, as the matrix high polymer for gelling such a carbonate, there may be used any one of high polymers having been generally used for a usual gel electrolyte. In particular, a fluorine based high polymer such as polyvinylidene fluoride, or vinylidene fluoride-hexafluoropropylene copolymer may be preferably used as the matrix high polymer from the viewpoint of stability of oxidation/reduction.

The high polymer solid electrolyte is composed of a lithium salt and a high polymer compound for dissolving the lithium salt. Examples of the high polymer compounds may include an ether based high polymer such as polyethylene oxide or a cross-linked high polymer thereof, a polymethacrylate ester based high polymer, an acrylate based high polymer, and a fluorine based high polymer such as polyvinylidene fluoride or vinylidene-hexafluoropropylene copolymer. These high polymers may be used singly or in combination. In particular, the fluorine based high polymer such as polyvinylidene fluoride or vinylidene-hexafluoropropylene copolymer may

be preferably used as the high polymer compound from the viewpoint of stability of oxidation/reduction.

As the lithium salt contained in the gel electrolyte or high polymer solid electrolyte, there can be used any one of the lithium salts having been generally used for a usual battery electrolyte. In particular, there may be used the lithium compounds (salts) such as lithium chloride, lithium bromide, lithium iodide, lithium chlorate, lithium perchlorate, lithium bromate, lithium iodate, lithium nitrate, lithium tetrafluoroborate, lithium hexafluorophosphate, lithium acetate, bis(trifluoromethanesulfonyl)imide,  $\text{LiAsF}_6$ ,  $\text{LiCF}_3\text{SO}_3$ ,  $\text{LiC}(\text{SO}_2\text{CF}_3)_3$ ,  $\text{LiAlCl}_4$ , and  $\text{LiSiF}_6$ . These lithium compounds may be used singly or in combination.

Of these lithium compounds, lithium hexafluorophosphate ( $\text{LiPF}_6$ ) and lithium tetrafluoroborate ( $\text{LiBF}_4$ ) may be preferably used from the viewpoint of stability of oxidation.

For the gel electrolyte, the lithium salt may be dissolved in the plasticizer at a concentration of 0.1 to 3.0 mol/l, preferably, 0.5 to 2.0 mol/l.

The components of the battery of the present invention, other than the above-described gel electrolyte containing a carbonate or the solid electrolyte may be



the same as those of a usual lithium ion battery.

For example, the negative electrode of the battery of the present invention may be made from a negative electrode material used for a usual lithium ion battery, typically, a carbon material such as a graphitization-retardant carbon material or a graphite based material in or from lithium can be doped or undoped. Examples of such carbon materials may include pyrocarbon, coke such as pitch coke, needle coke, or petroleum coke, graphite, vitreous carbon, a baked body of an organic high polymer compound (carbonization of phenol resin or furan resin by baking it at a suitable temperature), carbon fibers, and activated chacoal. As the material in or from which lithium can be doped or undoped, there may be also used polyacetylene, a high polymer such as polypyrrole or an oxide such as  $\text{SnO}_2$ . In formation of the negative electrode by using the above material, a known binder may be added thereto.

The positive electrode of the battery of the present invention may be configured by using a positive active material used for a usual lithium ion battery, typically a material selected from a metal oxide, a metal sulfide, or a specific high polymer in accordance with the kind of the battery. Examples of the positive active

materials may include a metal sulfide not containing lithium such as  $\text{TiS}_2$ ,  $\text{MoS}_2$ , or  $\text{NbSe}_2$ , a metal oxide not containing lithium such as  $\text{V}_2\text{O}_5$ , and a composite metal oxide containing lithium expressed by a chemical structure formula  $\text{Li}_x\text{MO}_2$  (M is one kind or more transition metals, and x is variable depending on the charging/discharging state of the battery, and is usually set in a range of 0.05 to 1.10). In the composite metal oxide  $\text{Li}_x\text{MO}_2$ , the transition metal may be selected from Co, Ni and Mn. Examples of the composite metal oxide  $\text{Li}_x\text{MO}_2$  may include  $\text{LiCoO}_2$ ,  $\text{LiNO}_2$ , and  $\text{LiNi}_y\text{Co}_{1-y}\text{O}_2$  ( $0 < y < 1$ ). The composite metal oxide  $\text{Li}_x\text{MO}_2$  can constitute a positive active material capable of generating a high voltage and enhancing the energy density. A plurality of these positive active materials may be combined to form the positive electrode. In formation of the positive electrode by using the above positive active material, a known conductive agent and a known binder may be used.

Electrode terminals (negative terminal lead 3 and positive electrode terminal lead 4) are connected to collectors of the negative and positive electrodes, respectively. The positive electrode may be made from a metal not melted at a high potential, such as aluminum, titanium, or an alloy thereof, and the negative electrode

may be made from copper, nickel, or an alloy thereof.

These electrode terminals are folded at winding terminal surfaces of the battery element 1 in such a manner as to be at the same level as that of the principal plane of the battery element 1 in order to improve the volume efficiency and to be easily led out of the battery element 1. The gas absorbable members 5 and 6 or the gas absorbable member 7 can effectively function from the viewpoint of prevention of short-circuit, and also function as a shock absorber.

The nonaqueous electrolyte battery may be used as a primary battery or secondary battery; however, it may be preferably used as a lithium ion secondary battery using a solid electrolyte or gel electrolyte.

The effects of the present invention will be more clearly understood by way of the following examples in which characteristics of sample batteries are variously evaluated.

<Example 1>

#### Production of Sample Battery

First, a negative electrode was produced as follows:

A negative electrode depolarizing mix was prepared by mixing 90 parts by weight of a pulverized powder of

graphite and 10 parts by weight of vinylidene fluoride-hexafluoropropylene copolymer. The negative electrode depolarizing mix was then diffused in N-methyl-2-pyrrolidone into slurry. The slurry was applied on one surface of a strip-like copper foil, as a negative electrode collector, having a thickness of 10  $\mu$ m, being dried, and compressed by a roll press, to produce a negative electrode.

A positive electrode was produced as follows:

First, a positive active material ( $\text{LiCoO}_2$ ) was obtained by mixing lithium carbonate and cobalt carbonate at a mol ratio of 0.5 : 1, and the resultant mixture was baked at 900°C for 5 hr. A positive electrode depolarizing mix was prepared by mixing 91 parts by weight of the positive active material ( $\text{LiCoO}_2$ ), 6 parts by weight of graphite as a conductive agent, and 10 parts by weight of vinylidene-hexafluoropropylene copolymer as a binder. The positive electrode depolarizing mix was then diffused in N-methyl-2-pyrrolidone into slurry. The slurry was applied on one surface of a strip-like aluminum foil, as a positive electrode collector, having a thickness of 20  $\mu$ m, being dried, and compressed by a roll press, to produce a positive electrode.

A positive electrode terminal lead and a negative

electrode terminal lead were welded on portions, where the active materials were not formed, of the positive electrode (aluminum foil) and the negative electrode (copper foil), respectively.

A gel electrolyte was obtained as follows:

First, a plasticizer was prepared by mixing 42.5 parts by weight of ethylene carbonate (EC), 42.5 parts by weight of propylene carbonate (PC), and 15 parts by weight of  $\text{LiPF}_6$ . An electrolyte solution was obtained by mixing and dissolving 30 parts by weight of the plasticizer and 10 parts by weight of vinylidene fluoride-hexafluoropropylene copolymer having a weight-average molecular weight  $M_w$  of 600,000 in 60 parts by weight of diethyl carbonate. The electrolyte solution was applied on the negative and positive electrodes, being impregnated therein, and left at ordinary temperature for 8 hr to evaporate the dimethyl carbonate, to thereby obtain a gel electrolyte.

The gel electrolyte sides of the negative and positive electrodes coated with the gel electrolyte were laminated to each other, being pressed and wound, to produce a winding type gel electrolyte battery element 1 (capacity: 550 mAh) having a size of  $3.3 \text{ cm} \times 5.2 \text{ cm} \times 3.3 \text{ mm}$  shown in Fig. 9.

Next, a hardening resin composition was obtained by mixing 30 parts by weight of NK Ester 1206PE (trade name, produced by Shin-Nakamura Chemical Co., Ltd.), 25 parts by weight of NK Ester 23G (trade name, produced by Shin-Nakamura Chemical Co., Ltd.), 40 parts by weight of carbon molecular sieve having an average particle size of 3  $\mu$ m (pulverized after vacuum heating and drying at 120°C for 24 hr), and 0.5 part by weight of 1173 (trade name, produced by Ciba-Geigy Limited) as a hardening agent, and vacuum degassing the resultant mixture. The hardening resin composition was irradiated with ultraviolet rays having an energy of 20 mJ/cm<sup>2</sup> for 1 min by using a high pressure mercury lamp to be solidified, to produce a plate-like gas absorbable member having a size of 3.3 cm $\times$  3.3 mm $\times$ 0.3 mm. In addition, the production of the above gas absorbable member was performed in a glove box filled with argon gas disposed in a dry room having a dew point of -45°C.

The battery element and the gas absorbable member thus produced were inserted in an outer covering member composed of a laminated film, and were sealed therein by heat-sealing the periphery of the outer covering member along a seal width of 5 mm for at 200°C for 10 sec by a heat sealer, to produce a sample battery.

<Example 2>

A sample battery was produced in the same manner as that in Example 1, except that diethyl carbonate was used in place of propylene carbonate constituting part of the plasticizer, and the content of the carbon molecular sieve having an average particle size of 3  $\mu\text{m}$  was changed from 40 parts by weight into 32 parts by weight.

<Example 3>

A sample battery was produced in the same manner as that in Example 1, except that 85 parts by weight of propylene carbonate was used in place of the mixed solvent of 42.5 parts by weight of ethylene carbonate and 42.5 parts by weight of propylene carbonate constituting parts of the plasticizer, and a pulverized powder of silica gel having an average particle size of 5  $\mu\text{m}$  was used in place of the carbon molecular sieve having an average particle size of 5  $\mu\text{m}$ .

<Example 4>

A sample battery was produced in the same manner as that in Example 1, except that a plate-like gel electrolyte battery element of a capacity of 1200 mAh having a size of 140 mm $\times$ 196 mm $\times$ 0.46 mm shown in Fig. 10 was produced in place of the winding type gel electrolyte battery element, and a laminated film having a gas

absorbable layer containing carbon molecular sieve having an average particle size of 3  $\mu\text{m}$  shown in Fig. 8 was used in place of the flat-like gas absorbable member.

<Example 5>

A sample battery was produced in the same manner as that in Example 4, except that diethyl carbonate was used in place of propylene carbonate constituting part of the plasticizer.

<Example 6>

A sample battery was produced in the same manner as that in Example 4, except that 85 parts by weight of propylene carbonate was used in place of the mixed solvent of 42.5 parts by weight of ethylene carbonate and 42.5 parts by weight of propylene carbonate constituting parts of the plasticizer, and a pulverized powder of silica gel having an average particle size of 5  $\mu\text{m}$  was used in place of the carbon molecular sieve having an average particle size of 3  $\mu\text{m}$ .

<Comparative Example 1>

A sample battery was produced in the same manner as that in Example 1, except that the carbon molecular sieve as the gas absorbable material was not used.

<Comparative Example 2>

A sample battery was produced in the same manner as



that in Example 2, except that the carbon molecular sieve as the gas absorbable material was not used.

<Comparative Example 3>

A sample battery was produced in the same manner as that in Example 3, except that the pulverized powder of silica gel as the gas absorbable material was not used.

<Comparative Example 4>

A sample battery was produced in the same manner as that in Example 4, except that the carbon molecular sieve as the gas absorbable material was not used.

<Comparative Example 5>

A sample battery was produced in the same manner as that in Example 5, except that the carbon molecular sieve as the gas absorbable material was not used.

<Comparative Example 6>

A sample battery was produced in the same manner as that in Example 6, except that the pulverized powder of silica gel as the gas absorbable material was not used.

Each of the sample batteries in Examples 1 to 6 and Comparative Example 1 to 6 was evaluated in terms of the ability of absorbing gas occurred in the battery element by battery reaction.

Each sample battery was initially charged at 5-hour rate of the theoretical capacity (1/5C), and over-charged

to 4.35 V at 2-hour rate of the theoretical capacity (1/2C). The sample battery in the over-charged state was left in a thermostat kept at 100°C for 100 hr.

After being left for 100 hr, the sample battery was taken out of the thermostat, and the swelling degree of the laminated film was visually observed at a time point directly after the sample was taken out of the thermostat and at a time point after the sample battery was cooled to ordinary temperature.

The criterion of evaluating the swelling degree of the laminated film was set as follows:

A sample battery, in which the laminated film is not swelled, the adhesion between the battery element and the laminated film is kept, and the battery shape is kept, is evaluated as a mark ○; a sample battery, in which the battery shape is substantially kept, but the battery element is separated from the laminated film whereby the adhesion is not kept, is evaluated as a mark △; and a sample battery, in which the laminated film is swelled and the battery shape is not kept, is evaluated as a mark ×.

The results of evaluating the gas absorbing abilities of the sample batteries in Examples 1 to 6 and Comparative Examples 1 to 6 are shown in Table 1.

Table 1

	Observation Result <sup>(1)</sup>	Observation Result <sup>(2)</sup>
Example 1	○	○
Example 2	△	○
Example 3	○	○
Example 4	○	○
Example 5	△	○
Example 6	○	○
Comparative Example 1	×	×
Comparative Example 2	×	×
Comparative Example 3	×	×
Comparative Example 4	×	×
Comparative Example 5	×	×
Comparative Example 6	×	×

Observation Result <sup>(1)</sup>: directly after the sample battery is taken out of the thermostat

Observation Result <sup>(2)</sup>: after the sample battery is cooled to ordinary temperature

As is apparent from Table 1, in each of the sample batteries in Examples 2 and 5, the laminated film is loosened directly after the sample battery is taken out of the thermostat kept at 100°C; however, in each of the sample batteries in Examples 1 to 6, in the state after the sample battery is cooled to ordinary temperature, the laminated film is not loosened, the adhesion between the battery element and the laminated film is kept, and the battery shape is kept.

On the contrary, in each of the sample batteries in

Comparative Examples 1 to 6, the laminated film is swelled and the battery shape is not kept directly after the sample battery is taken out of the thermostat kept at 100°C, and the swelling of the laminated film is left after the sample battery is cooled to ordinary temperature.

As a result, according to the nonaqueous electrolyte battery, since the gas absorbable material is inserted between the battery element and the laminated film or contained in the laminated film, even if the battery undergoes an abnormal environmental change such as overcharging or an abnormal temperature rise, gas occurred in the battery element is absorbed in the gas absorbable material, with a result that a change in shape of the battery due to the swelling of the laminated film can be suppressed.

While the preferred embodiments of the present invention have been described using the specific terms, such description is for illustrative purposes only, and it is to be understood that changes and variations may be made without departing from the spirit or scope of the following claims.

What is Claimed is:

1. A nonaqueous electrolyte battery comprising:  
a battery element contained in an outer covering member composed of a laminated film and sealed therein by heat seal; and

a gas absorbable material interposed between the outermost layer of said outer covering member and said battery element.

2. A nonaqueous electrolyte battery according to claim 1, wherein said gas absorbable material is a porous metal oxide or a porous carbon material

3. A nonaqueous electrolyte battery according to claim 2, wherein said porous metal oxide is one kind selected from a group consisting of zeolite, alumina, molecular sieve, titania, silica gel, and zirconia.

4. A nonaqueous electrolyte battery according to claim 2, wherein said porous carbon material is activated carbon or carbon molecular sieve.

5. A nonaqueous electrolyte battery according to claim 1, wherein said gas absorbable material is mixed with a resin material and the mixture is molded to form a gas absorbable member, and said gas absorbable member is inserted between the outermost layer of said outer covering member and at least one or more planes of said

battery element.

6. A nonaqueous electrolyte battery according to claim 1, wherein said gas absorbable material is contained in said laminated film.

7. A nonaqueous electrolyte battery according to claim 1, wherein an electrolyte constituting part of said battery element is a gel electrolyte or solid electrolyte containing a matrix high polymer and a lithium salt.

8. A nonaqueous electrolyte battery according to claim 1, wherein a negative electrode constituting part of said battery element contains a material in or from which lithium is allowed to be doped or undoped.

9. A nonaqueous electrolyte battery according to claim 8, wherein said material in or from which lithium is allowed to be doped or undoped is a carbon material.

10. A nonaqueous electrolyte battery according to claim 1, wherein a positive electrode constituting part of said battery element contains a composite oxide of lithium and a transition metal.

11. A nonaqueous electrolyte battery according to claim 1, wherein said battery is a secondary battery.

# ABSTRACT OF THE DISCLOSURE

Disclosed is a nonaqueous electrolyte battery including a battery element contained in an outer covering member composed of a laminated film and sealed therein by heat seal, and a gas absorbable material interposed between the outermost layer of the outer covering member and the battery element. Such a battery is advantageous in that even if gas occurs in the battery element, the laminated film as the outer covering member is prevented from being swelled.

FIG. 1

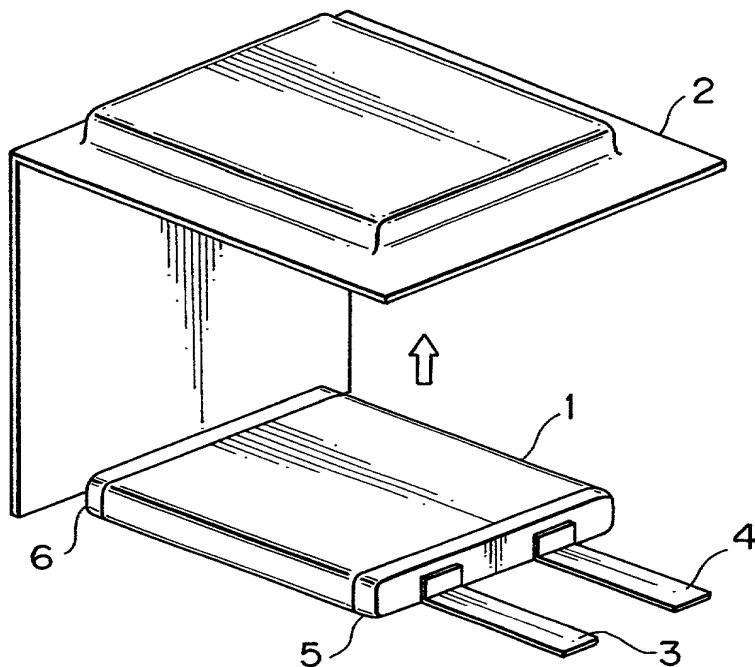


FIG. 2

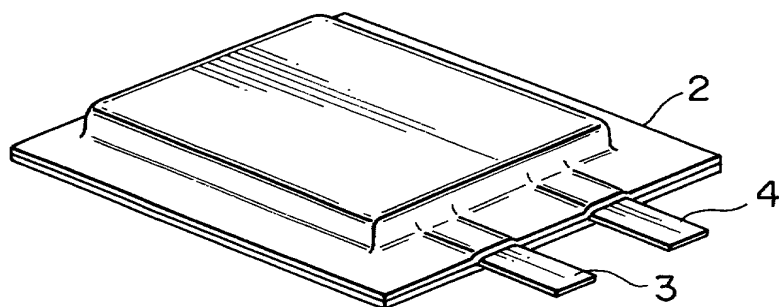




FIG. 3

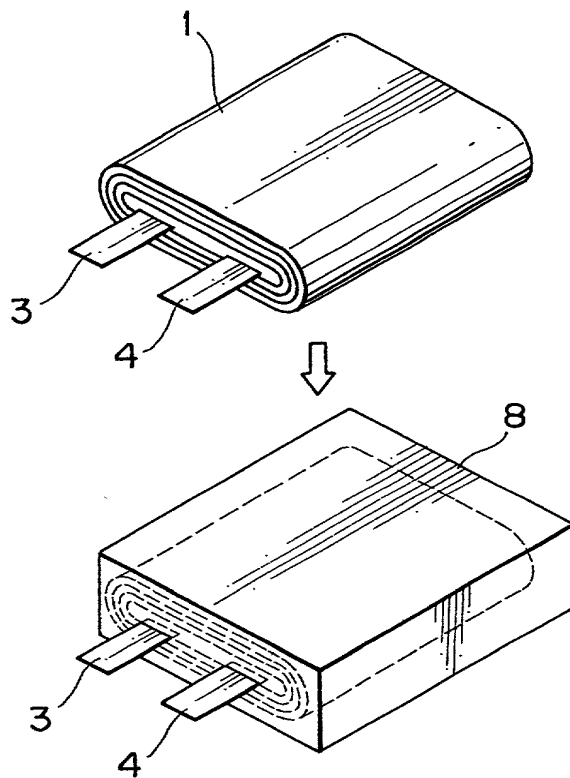


FIG. 4

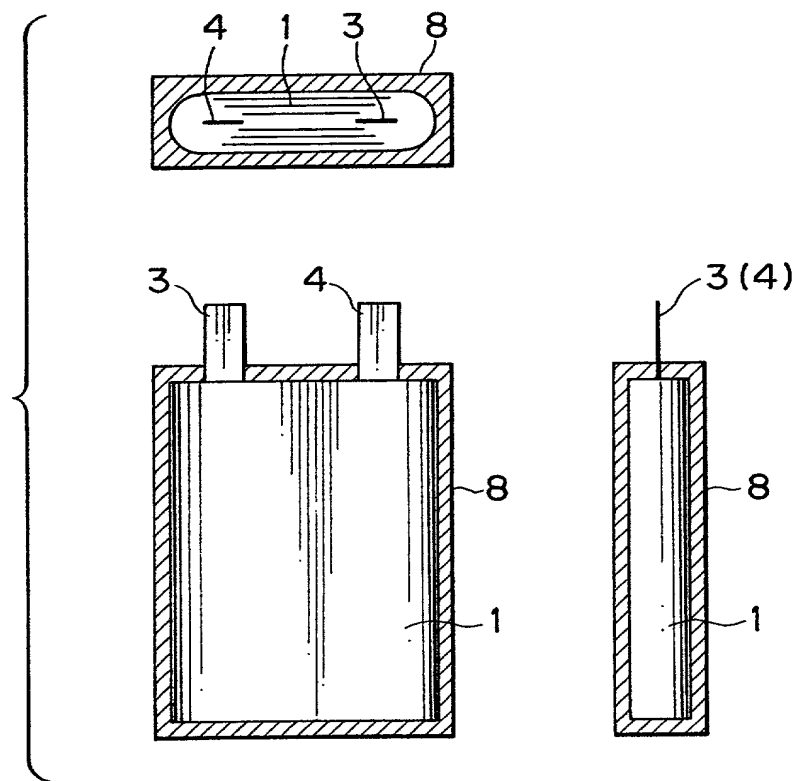


FIG. 5

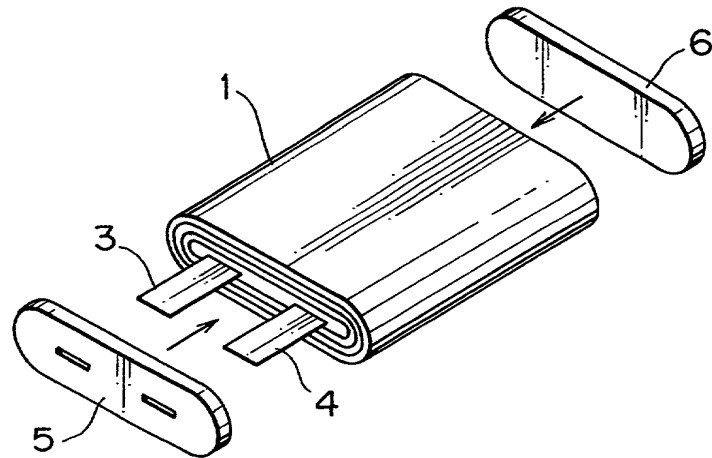


FIG. 6

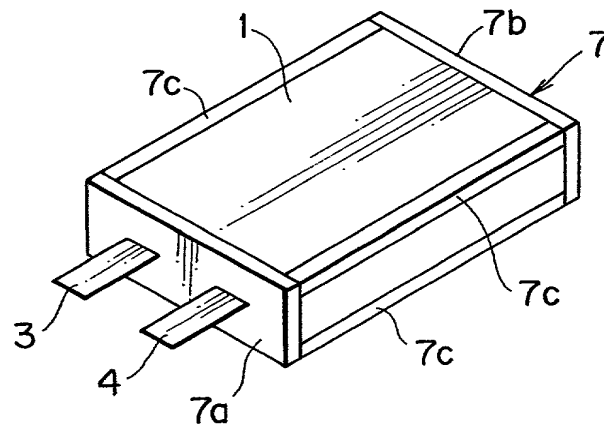


FIG. 7

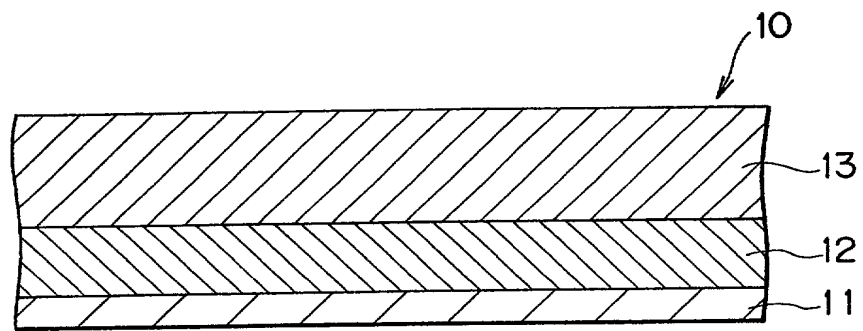


FIG. 8

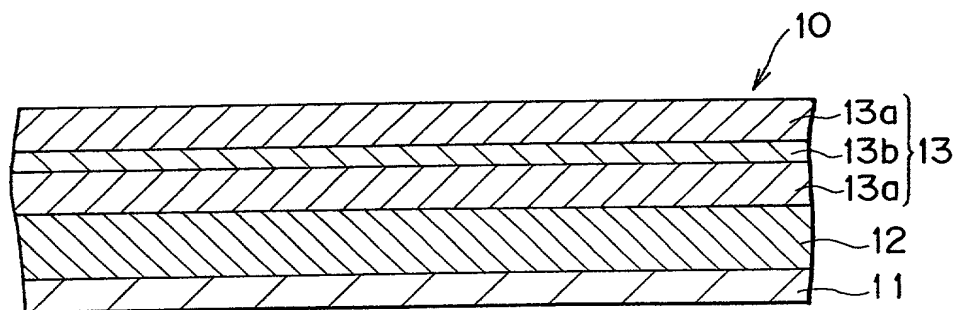


FIG. 9

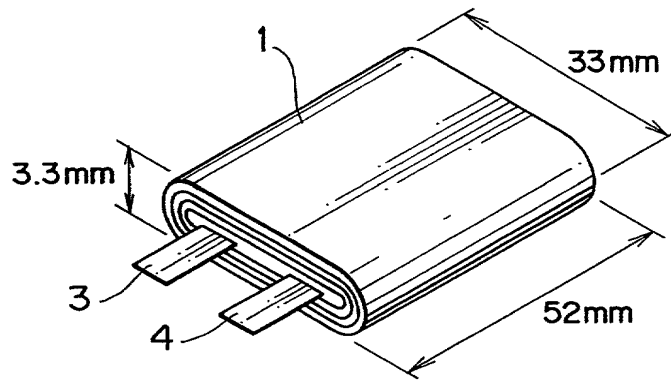
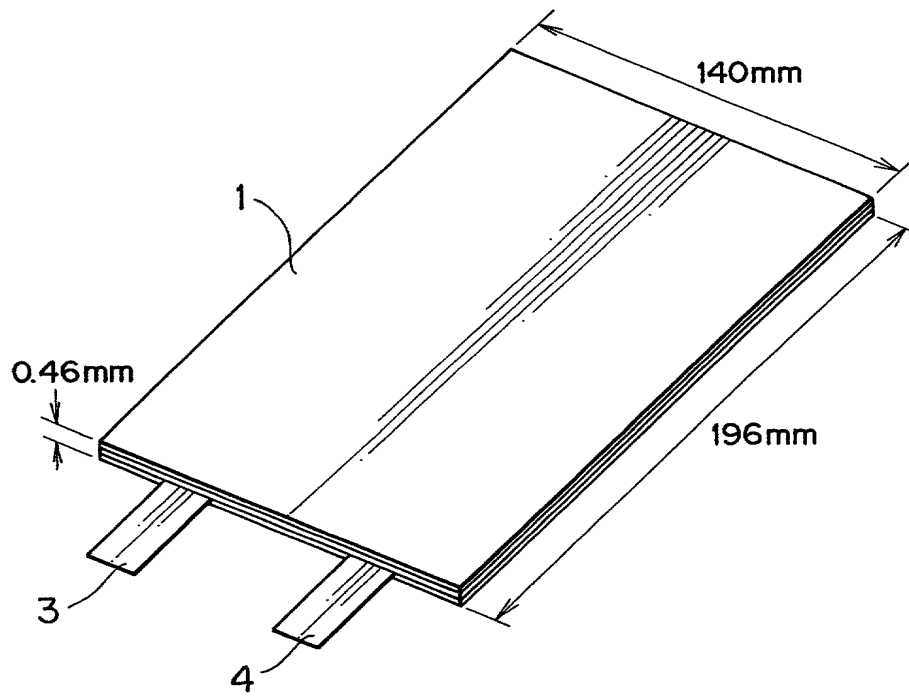


FIG. 10



# DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

## 特許出願宣言書及び委任状

### Japanese Language Declaration

### 日本語宣言書

下記の氏名の発明者として、私は以下の通り宣言します。

私の住所、私書箱、国籍は下記の私の氏名の後に記載された通りです。

下記の名称の発明に関して請求範囲に記載され、特許出願している発明内容について、私が最初かつ唯一の発明者（下記の氏名が一つの場合）もしくは最初かつ共同発明者であると（下記の名称が複数の場合）信じています。

上記発明の明細書（下記の欄でx印がついていない場合は、本書に添付）は、

☐ \_\_月\_\_日に提出され、米国出願番号または特許協定条約国際出願番号を\_\_\_\_とし、  
（該当する場合）\_\_\_\_に訂正されました。

私は、特許請求範囲を含む上記訂正後の明細書を検討し、内容を理解していることをここに表明します。

私は、連邦規則法典第37編第1条56項に定義されたとおり、特許資格の有無について重要な情報を開示する義務があることを認めます。

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled

**“NONAQUEOUS ELECTROLYTE BATTERY”**

the specification of which is attached hereto unless the following box is checked:

☐ was filed on \_\_\_\_\_ as United States Application Number or PCT International Application Number \_\_\_\_\_ and was amended on \_\_\_\_\_ (if applicable)

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, Section 1.56.

私は、米国法典第35編119条(a)-(d)項又は365条(b)項に基づき下記の、米国以外の国の少なくとも一カ国を指定している特許協力条約365(a)項に基づき国際出願、又は外国での特許出願もしくは発明者証の出願についての外国優先権をここに主張するとともに、優先権を主張している、本出願の前に出願された特許または発明者証の外国出願を以下に、枠内をマークすることで、示しています。

Prior Foreign Application(s)

外国での先行出願

P11-341373 Japan November 30, 1999

(Number) (番号)	(Country) (国名)	(Day Month Year Filed) (出願年月日)
------------------	-------------------	-----------------------------------

(Number) (番号)	(Country) (国名)	(Day Month Year Filed) (出願年月日)
------------------	-------------------	-----------------------------------

I hereby claim foreign priority under Title 35, United States Code, Section 119(a)-(d) or 365(b) of any foreign application(s) for patent or Inventor's certificate or 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below, by checking the box, any foreign application for patent or Inventor's certificate or PCT International application having a filing date before that of the application on which priority is claimed:

Priority Not Claimed

優先権主張なし

(Number)	(Country)	(Day Month Year Filed)
----------	-----------	------------------------

(Number)	(Country)	(Day Month Year Filed)
----------	-----------	------------------------

私は、第35編米国法典119条(e)項に基づいて下記の米国特許出願規定に記載された権利をここに主張いたします。

(Application No.) (出願番号)	(Filing Date) (出願日)
-----------------------------	------------------------

I hereby claim the benefit under Title 35, United States Code, Section 119(e) of any United States provisional application(s) listed below.

(Application No.) (出願番号)	(Filing Date) (出願日)
-----------------------------	------------------------

私は、下記の米国法典第35編120条に基づいて下記の米国特許出願に記載された権利、又は米国を指定している特許協力条約365条(c)に基づき権利をここに主張します。また、本出願の各請求範囲の内容が米国法典第35編112条第1項又は特許協力条約で規定された方法で先行する米国特許出願に開示されていない限り、その先行米国出願書提出日以降で本出願書の日本国内または特許協力条約国際提出日までの期間中に入手された、連邦規則法典第37編1条56項で定義された特許資格の有無に関する重要な情報について開示義務があることを認識しています。

I hereby claim the benefit under Title 35, United States Code, Section 120 of any United States application(s) or 365(c) of an PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of Title 35, United States Code, Section 112, I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, Section 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application.

(Application No.)  
(出願番号)

(Filing Date)  
(出願日)

(Status: patented, pending, abandoned)  
(現況: 特許許可済、係属中、放棄済)

(Application No.)  
(出願番号)

(Filing Date)  
(出願日)

(Status: patented, pending, abandoned)  
(現況: 特許許可済、係属中、放棄済)

私は、私自身の知識に基づいて本宣言書中で私が行なう表明が真実であり、かつ私の入手した情報と私の信じることに基づく表明が全て真実であると信じていること、さらに故意になされた虚偽の表明及びそれと同等の行為は米国法典第18編第1001条に基づき、罰金または拘禁、もしくはその両方により処罰されること、そしてそのような故意による虚偽の声明を行なえば、出願した、又は既に許可された特許の有効性が失われることを認識し、よってここに上記のごとく宣誓を致します。

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

委任状: 私は下記の発明者として、本出願に関する一切の手続きを米特許商標局に対して遂行する弁理士または代理人として、下記の者を指名いたします。(弁理士、または代理人の氏名及び登録番号を明記のこと)

POWER OF ATTORNEY: As a named Inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (list name and registration number)

David R. Metzger (Reg. No. 32,919), Joseph A. Mahoney (Reg. No. 38,956), Howard B. Rockman (Reg. No. 22,190), Jordan A. Sigale, (Reg. No. 39,028), Michael A. Molano (Reg. No. 39,777), Michael L. Kiklis (Reg. No. 38,939), Janelle D. Strode (Reg. No. 34,738), Kevin W. Guynn (Reg. No. 29,972), Jennifer Hammond (Reg. No. 41,814), Lana Knedlik (Reg. No. 42,748), John F. Griffith (Reg. No. 44,137), Marina Saito (Reg. No. 42,121), Alison P. Schwartz (Reg. No. 43,863), Christopher P. Rauch (Reg. No. 45,034), Francisco Rubio-Campos (Reg. No. 45,358), Brian J. Gill (Reg. No. 46,727) and Shashank S. Upadbye, all members of the firm of Sonnenschein, Nath & Rosenthal

Send Correspondence to:

書類送付先

David R. Metzger  
Sonnenschein Nath & Rosenthal  
P.O. Box #061080  
Wacker Drive Station  
Chicago, Illinois 60606-1080

直接電話連絡先: (名前及び電話番号)

Direct Telephone Calls to: (name and telephone number)

312/876-2578



唯一または第一発明者名	Full name of sole or first inventor: TSUYONOBU HATAZAWA
発明者の署名 日付	Inventor's signature Date
住所	Residence Miyagi, Japan
国籍	Citizenship Japan
私書箱	Post Office Address c/o Sony Corporation 7-35, Kitashinagawa 6-chome
	Shinagawa-ku, Tokyo 141, Japan

(第三以降の共同発明者についても同様に記載し、署名をすること)

第二共同発明者	Full name of second joint inventor, if any: TOMITARO HARA
発明者の署名 日付	Inventor's signature Date
住所	Residence Miyagi, Japan
国籍	Citizenship Japan
私書箱	Post Office Address c/o Sony Corporation 7-35, Kitashinagawa 6-chome
	Shinagawa-ku, Tokyo 141, Japan
第二共同発明者	Full name of third joint inventor, if any: TAKAHIRO ENDO
発明者の署名 日付	Inventor's signature Date
住所	Residence Miyagi, Japan
国籍	Citizenship Japan
私書箱	Post Office Address c/o Sony Corporation 7-35, Kitashinagawa 6-chome
	Shinagawa-ku, Tokyo 141, Japan

## Japanese Language Declaration

(日本語宣言書)

第二共同発明者		Full name of fourth joint inventor, if any: KAZUHITO HATTA	
発明者の署名	日付	Inventor's signature	Date
住所		Residence Miyagi, Japan	
国籍		Citizenship Japan	
私書箱		Post Office c/o Sony Corporation 7-35, Kitashinagawa 6-chome	
		Shinagawa-ku, Tokyo 141, Japan	
第二共同発明者		Full name of fifth joint inventor, if any:	
発明者の署名	日付	Inventor's signature	Date
住所		Residence	
国籍		Citizenship	
私書箱		Post Office Address	
第二共同発明者		Full name of sixth joint inventor, if any:	
発明者の署名	日付	Inventor's signature	Date
住所		Residence	
国籍		Citizenship	
私書箱		Post Office Address	
第二共同発明者		Full name of seventh joint inventor, if any:	
発明者の署名	日付	Inventor's signature	Date
住所		Residence	
国籍		Citizenship	
私書箱		Post Office Address	